PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5: (11) International Publication Number: WO 90/12081 A1 C11D 3/395, 17/00, 3/37 (43) International Publication Date: 18 October 1990 (18.10.90)

(21) International Application Number: PCT/US89/03313 (74) Agents: BYRNE, Linda, M. et al.; 1000 Norwest Center,

(22) International Filing Date: I August 1989 (01.08.89)

31 March 1989 (31.03.89) 331,695 US

(71) Applicant: ECOLAB INC. [US/US]; 370 North Wabasha Street, St. Paul, MN 55102 (US).

(72) Inventors: BULL, Sandra, L.; 1254 Wilderness Curve, Eagan, MN 55123 (US). GLADFELTER, Elizabeth, J.; 1574 Arona Street. Falcon Heights, MN 55113 (US). OL-SON, Keith, E.: 13952 Eveleth Court, Apple Valley, MN 55124 (US).

55 East Fifth Street, St. Paul, MN 55101 (US).

(81) Designated States: AT (European patent), AU, BE (Euro-+ pean patent), CH (European patent), DE (European patent), FR (European patent), GB (European patent), IT (European patent), JP, LU (European patent), NL (European patent), SE (European patent).

Published

With international search report.

(54) Title: CAST DETERSIVE SYSTEMS

(57) Abstract

(30) Priority data:

A general purpose detersive system and detersive system useful in warewashing and in laundry processes have been developed in which highly active encapsulated halogen sources have been incorporated into cast, solid detersive systems containing oxidizable organics that are highly reactive with the active chlorine sources. The stability of the detersive systems thus formulated has been shown to be sufficient to permit the cast materials to be storage stable for a sufficient period to permit the manufacture, distribution, sale and consumption of the cast materials before the availability of either the organic materials or the active halogen drops below an effective level.

DESIGNATIONS OF "DE"

Until further notice, any designation of "DE" in any international application whose international filing date is prior to October 3, 1990, shall have effect in the territory of the Federal Republic of Germany with the exception of the territory of the former German Democratic Republic.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT AU BB BE BF BG BJ CA CF CG CH DE DK	Austria Austrialia Barbados Belgium Burkina Fasso Bulgaria Benin Brazil Canada Centrial African Republic Congo Switzerland Cameroon Germany, Federal Republic of Denmark	ES FI FR GA GB HU IT JP KP KR U LK U MC	Spain Finland France Gabon United Kingdom Hungary Italy Japan Democratic People's Republic of Korea Republic of Korea Liechtenstein Sri Lanka Luxembourg Monaco	MG ML MR MP NL NO SO SE SN J P T G	Madagascar Mali Mauritania Malawi Netherlands Norway Romania Sudan Sweden Senegal Soviet Union Chad Togo United States of America

WO 90/12081 - 1 - PCT/US89/03313

CAST DETERSIVE SYSTEMS .

Field of the Invention

The invention relates to a detersive system containing 5 soil removing detergents, an encapsulated halogen source and an organic component such as a polyalkylene oxide polymer, and to the use of the system in cleaning. specifically the detersive systems of the invention can contain an active encapsulated halogen source and organic 10 component, preferably an alkylene oxide polymer, which are co-stable at relatively high concentration during manufacture, storage, distribution, sale and use. Such detersive systems can be used in a variety of environments including general purpose cleaning, laundry, warewashing, 15 etc., while producing an effective concentration of active chlorine for stain removal and sanitizing purposes in the presence of organic materials such as defoamers, soil removing nonionic surfactants and other polyalkylene oxide polymers.

20 <u>Background of the Invention</u>

Detersive systems have been used for many years in cleaning environments such as laundry, warewashing, hard surface cleaning and other general cleaning applications. Typically detersive systems are concentrates comprising mixtures of cleaning ingredients that, just prior to use, are mixed with water to form a cleaning medium or final use composition.

Many detersive systems contain organic materials that function in a variety of modes. Such organics can act as 30 defoaming agents, dispersing agents, soil removing surfactants, thickening agents, casting aids, antiredeposition aids, viscosity modifiers, brightening agents, One important class of organic materials polyalkylene oxide polymers. The polyalkylene oxide polymers derive their properties from the presence of large polymeric units derived from ethylene oxide, propylene oxide, heteric polymers thereof or block copolymers

thereof.

In a variety of use environments for such detersive systems, the presence of oxidizing available halogen such as chlorine or bromine can be important. Such halogen 5 compounds can clean and remove certain residues, and can remove stains by oxidizing and destroying color generating functional groups in organic molecules. presence of sufficient concentrations of active chlorine can kill microorganisms and provide an antimicrobial or 10 sanitizing action. However, detersive manufacturers have been reluctant to produce detersive systems in which organic materials such as polyalkylene oxide polymers and active chlorine yielding substances come into direct contact. Such contact can have undesirable 15 safety and performance consequences.

In the manufacture of a variety of detersive systems, contacting many organic substances with active chlorine compounds is hazardous. At elevated temperatures or at high concentrations, contacting such organic compounds with 20 active chlorine-yielding substances can result in a rapid reaction between the chlorine substances and the organic materials resulting in the production of large volumes of smoke or fire. If the detersive systems are successfully manufactured without rapid reaction between the chlorine 25 and polyalkylene oxide components, the materials during manufacture, packaging, storage, distribution, sale and use can slowly react causing a significant depletion in the concentration of both the organic materials and the active chlorine substance. Such reactions commonly occur at a 30 rate such that the concentration of active chlorine in the detersive system is below the concentration necessary for sanitizing properties and even is often below concentration required for stain removal. Further such reactions can reduce the concentration of the organic 35 materials to a level that results in substantially reducing the level of properties derived from the polymer.

We have found that active halogen can be lost through

WO 90/12081 – 3 – PCT/US89/03313

a spontaneous degradation of active chlorine and by reactivity of chlorine with functional groups commonly found in organics used in detersive systems. The common functional groups reactive with active chlorine compounds include olefinic bonds commonly found in unsaturated fatty acids which are a common composition of alkali metal soaps, hydroxyl groups typically found in organic builders, amine groups, ether groups common in polyalkylene oxide polymers and aromatic rings typically found in alkyl bezzene sulfonates, alkyl phenol ethoxylates, etc. The prior art commonly kept the concentration of organics less than 5 wt-% to reduce the harmful effects of the organic/chlorine interaction.

Accordingly a substantial need exists in the art for detersive systems containing significant effective concentrations (greater than 5 wt-%) of organic materials in the form of surfactants, foam suppressing agents and other functional materials, and containing effective concentrations (greater than 5 wt-%) of active chlorine from chlorine yielding compounds.

Brief Description of the Invention

We have found that a cast, solid detersive system containing an effective amount of a source of alkalinity, an effective hardness sequestering agent, at least 5 wt-% 25 of an encapsulated source of chlorine and at least 5 wt-% of an organic such as polyalkylene oxide polymer is an effective detersive system that can be used in general purpose cleaning, laundry, warewashing, hard cleaning and sanitizing, and a variety of other end uses. 30 We have surprisingly found that such materials when cast in a solid form result in substantial stability of effective concentration of both chlorine and organic materials during manufacture, storage, distribution and We have found that for periods typical in the 35 lifetime of such a product, the products can maintain a substantial concentration of active chlorine effective concentration of organics until the product is

entirely consumed.

Detailed Discussion of the Invention

The detersive systems of the invention can comprise an effective amount of a source of alkalinity, an effective amount of a hardness sequestering agent, at least 5 wt-%, up to about 50 wt-%, of an encapsulated source of chlorine compound, and at least 5 wt-%, up to about 50 wt-%, of an organic composition such as a polyalkylene oxide polymer. The detersive systems of this invention are typically in the form of a cast, solid material wherein the material is in the form of a large, solid mass having a minimum weight of about 50 grams held within a disposable container. The detersive system or concentrate can be dispensed from the container or can be removed from its container and placed into the dispenser for dilution and use at the use site.

Inorganic Source of Alkalinity

The detersive systems of the invention can contain an inorganic source of alkalinity. Sources of alkalinity identified as being useful in combination with the other 20 components of the detersive systems of the invention include but are not limited to the following: alkali metal hydroxide, alkali metal phosphate, alkali metal carbonate, alkali metal bicarbonate, alkali metal sequicarbonate, alkali metal borate, alkali metal silicate, and mixtures 25 thereof. Alkali metal hydroxides are typically used where the detersive system requires strong cleaning action and Silicates (M20:SiO2 compounds having a ratio highest pH. from about 2:1 to 1:3.6 wherein M is an alkali metal ion) are typically a reaction product between an alkali metal 30 hydroxide and silica. Such silicates are primarily used as a source of alkalinity where moderate strength and pH are required. Carbonate and borate sources of alkalinity are typically used in compositions having an alkaline pH but do not require the strength of alkalinity of silicates or 35 hydroxides.

Hardness Sequestering Agents

The detersive systems of the invention typically

WO 90/12081 - 5 - PCT/US89/03313

contain a hardness sequestering agent to reduce the harmful effects of divalent and trivalent metal ions on the components of the detersive systems. The detersive systems of the invention can contain both organic and inorganic hardness sequestering agents. Such sequestering agents are typically in the form of polyanionic materials.

Inorganic hardness sequestering agents include alkali metal condensed phosphates in the form of pyrophosphate, tripolyphosphate, hexametaphosphate, etc. Organic hardness sequestering agents include polymeric and copolymeric compositions having pendent carboxylic acid functionality derived from carboxylic acid containing monomers such as acrylic acid, methacrylic acid, maleic acid, itaconic acid, fumaric acid and mixtures thereof, etc.; alkali metal salts of organic substituted phosphonic acid and polyphosphonic acids, alkali metal salts of gluconic acid, alkali metal salts of ethylene diamine tetraacetic acid, alkali metal salts of nitrilotriacetic acid, and mixtures thereof.

Organic Materials

A variety of organic functional materials can be used 20 in the detersive systems of the invention. Such functional materials include absorbents, adsorbents, antimicrobials, antioxidants, anti-soil agents, perfumes, dyes, binders, chelating agents, corrosion inhibitors, coupling agents, 25 defoamers, dispersants, solubilizers, stabilizers, thickeners, and UV absorbers. Examples of such absorbents adsorbents are microcrystalline cellulose, ricinoleate, free-flowing malto dextrin, powdered acrylate copolymers, and others. Anti-soil or anti-soil 30 redeposition agents that can be used include fatty acid amides, fluorocarbon surfactants, complex phosphate esters, styrene maleic anhydride copolymers, and cellulosic derivatives such as hydroxyethyl cellulose, hydroxypropyl cellulose, and others. Powder binders that can be used to 35 aid in the formation οĒ solid materials microcrystalline cellulose, long chain lactate esters, long chain oleate esters, polyacrylamides, microcrystalline

waxes, polyvinyl alcohol resins, polyethylene, polyvinyl pyrrolidone, and others.

Defoamers that can be used in the detersive systems of the invention include high molecular weight C_{10-40} linear primary alcohols, polyalkylene glycols, well known silicone defoamers, certain acrylate copolymers, and others.

One important class of functional organic materials in the manufacture of detersive systems of the invention comprises organic surfactants. A wide variety of surfactants can be used in the detersive systems of the invention including anionic surfactants, zwitterionic surfactants (containing both anionic and cationic groups), cationic surfactants and nonionic surfactants. Anionic surfactants include alkyl carboxylate (sodium and potassium soaps), alkyl sulfate, alkyl ether sulfate, alkyl benzene sulfonate, alkyl sulfonate, sulfonated fatty acid ester, sulfosuccinate surfactant.

Nonionic surfactants typically in the form of compositions having polyalkylene oxide polymers as a 20 portion of the surfactant molecule can be useful in the detersive systems of the invention.

Nonionic surfactants which are advanta; eously employed in the composition of this invention include, but are not to, the following polyoxyalkylene nonionic limited detergents: $C_8 - C_{22}$ normal fatty alcohol-ethylene oxide or propylene oxide condensates, i.e., condensation products of one mole of a fatty alcohol containing 8 to 22 carbon atoms with from 2 to 20 moles of ethylene oxide or propylene oxide; polyoxypropylene-polyoxyethylene condensates having 30 the formula $\mathrm{HO(C_2H_4O)_x(C_3H_6O)_y}$ -H wherein $\mathrm{(C_2H_4O)_x}$ equals at least 15% and $(C_3H_6O)_v$ equals 20-90% of the total weight of alkyl polyoxypropylenepolyoxyethylene compound; condensates having the formula RO-(C3 $^{\rm H}_6$ O) $_{\rm x}$ (C2 $^{\rm H}_4$ O) $_{\rm y}$ H where R is a C_1-C_{15} alkyl group and x and y each represent an integer of from 2 to 98; polyoxyalkylene glycols as described in U.S. Pat. No. 3,048,548; butylene oxide capped alcohol ethoxylates having the formula $R(OC_2H_4)_v(OC_4H_9)_vOH$ WO 90/12081 - 7 - PCT/US89/03313

where R is a C_8 - C_{18} alkyl group and y is from about 3.5 to 10 and x is an integer from about 0.5 to 1.5; benzyl ethers of polyoxyethylene condensates of alkyl phenols having the formula

5

where R is a $C_6^{-C}_{20}$ alkyl group and x is an integer of from 5 to 40; and alkyl phenoxy polyoxyethylene ethanols having the formula

$$R \left(OC_{2}H_{4}\right)_{x}OH$$

where R is a C_8 - C_{20} alkyl group and x is an integer of from 3 to 20.

20 Non-Surfactant Polyoxyalkylene Groups

Nonsurfactant polyalkylene oxide polymers are used in the manufacture of the detersive systems of the invention as a binder or as a casting agent in which the materials of the invention are blended with the casting agent at elevated temperatures and the cooling of the blended material solidifies the casting agent resulting in a stable cast solid. Such polyalkylene oxide polymers comprise homopolymers of ethylene oxide (EO), homopolymers of propylene oxide (PO), heteric EO/PO copolymers or block EO/PO copolymers, for example, those of molecular weight from 1,000 to 10,000, especially about 6,000 to 8,000. These materials are not to be considered as surfactant materials in the context of the present invention.

Additional Ingredients

The detersive systems of the invention can contain anti-foaming agents, typically in an amount of from about 0.001% to about 2%, preferably 0.05 to 1%. Such foam

WO 90/12081 PCT/US89/03313

suppressing or foam regulating agents include silicone compounds, phosphate esters, microcrystalline slack waxes, etc. In somewhat greater detail, preferred suds suppressing agents are silicone materials which are siloxanes having the formula

10

35

wherein X is from about 20 to about 2,000 and R and R' are each independently selected from the group consisting of C₁₋₂₀ alkyl or alkyl substituted aryl groups. Preferred groups include methyl, ethyl, propyl, butyl, and phenyl. Polydimethyl siloxanes having a molecular weight in the range of 200 to 200,000 and higher are useful suds controlling agents. Other suds suppressing agents include alkyl phosphate esters such as monostearyl phosphate and microcrystalline waxes having a melting point from about 65° C. to about 100° C. and having a molecular weight from 400 to about 1,000.

The detersive systems of the invention can contain 25 neutral organic and inorganic fillers such as sodium sulfate and sodium chloride. Organic fillers that can be used in the invention include starch, sugars, alk:lene 1 glycols having from to 10 carbon atoms, Additionally the compositions can contain proteolytic and 30 amylolytic enzymes, tarnish inhibitors such as benzotriazol, antibacterial agents, anti-soil redeposition agents, soil suspending agents, dyes, perfumes, silicate, glass and aluminum tarnish suppressing agents.

Method of Casting the Detersive Systems

The components of the detersive systems are typically cast in a disposable container which also acts as a mold. The detersive system can be dispensed from the container or can be removed from the container prior to insertion in the

WO 90/12081 – 9 – PCT/US89/03313

dispenser. Alternatively the detersive system components can be cast in a reusable mold wherein the cast material is removed from the mold and placed into a separate disposable container for shipment and use.

The detersive system can be cast in virtually any order of addition of materials to a mixing unit. Once uniform the material for casting is then placed in an appropriate mold.

A preferred mode of casting the material is to place
the low molecular weight and water soluble materials into a
mixing chamber in conjunction with any water used in the
manufacture of the materials. Next the higher molecular
weight or insoluble materials are slowly added to the
contents of the mixing unit until a uniform suspension is
obtained. Lastly the active halogen source can be added to
the mixture. The mixture is carefully agitated at a slow
rate such that the integrity of the encapsulate is not
compromised.

Encapsulate

The encapsulated sources of halogen of this invention comprise a core of an active halogen compound and at least one coating layer. Preferably the encapsulated sources of halogen have a core and two or more coating layers. If one layer is used it preferably comprises an inorganic coating of a composition compatible with the halogen source, however certain coatings of a synthetic detergent an be used. If two layers are used the first layer comprises a typically inert or inorganic coating agent and the slond layer comprises an organic layer or a synthetic detergent layer.

<u>Halogen Source</u>

The halogen releasing substances suitable as a core material include halogen components capable of liberating active halogen species such as a free elemental halogen or -OX- wherein X is Cl or Br, under conditions normally used in detergent-bleaching cleaning processes. Preferably the halogen releasing compound releases chlorine or bromine

species. The most preferred halogen releasing compound releases chlorine. Chlorine releasing compounds include potassium dichloroisocyanurate, sodium dichloroisocyanurate, chlorinated trisodium phosphate, calcium hypo-5 chlorite, lithium hypochlorite, monochloroamine, dichloropentaisocyanurate, 1,3-dichloro-5,5-dimethvl hydantoin, paratoluene sulfondichloro-amide, trichloromelamine, N-chloromelamine, N-chlorosuccinimide, N,N'-dichloroazodi-carbonamide, N-chloro-acetyl-urea, N,Nchlorinated 10 dichlorobiuret, dicyandiamide, trichlorocya suric acid, and dichloroglycoluril.

Sodium dichloroisocyanurate, potassium dichloroisocyanurate and the dihydrates thereof are the most preferred oxidizing chlorine source suitable as a core substance.

15 These materials are commercially available and may be obtained from sources such as Monsanto or Olin Corp.

Coating Materials

The coating on the active halogen source can be a single or multiple layer coating. Single layer coatings can comprise virtually any inert organic or inorganic coating material stable with a halogen source that is solid at room temperature. In the instance that a multiple layer coating is used, often the first layer is an inorganic layer where the second layer comprises an organic layer that can be drawn from a variety of sources.

Nearly any substance may be employed as the first coating so long as it is substantially halogen inert and is a solid at normal storage temperatures typically between 30° and 100° F. (~1° to 38° C.). The coating material is preferably inert with respect to the core material. If the inner co ting material is potentially reactive with the core material, the core material may be initially coated with an inert material to prevent or retard any reaction between the core and this first coat, the initial coat acting as a chemical parrie between the core and the first and other Payers. Useful inorganics in the coating material include alkilies such as acdium carbonate, sodium

WO 90/12081 - 11 - PCT/US89/03313

bicarbonate, sodium sequicarbonate, sodium borate, potassium bicarbonate, potassium sequicarbonate, potassium phosphates such as diammonium monocalcium phosphate, tricalcium phosphate, calcium 5 pyrophosphate, iron pyrophosphate, magnesium phosphate, monopotassium orthophosphate, potassium pyrophosphate, disodium orthophosphate, trisodium orthophosphate, tetrasodium pyrophosphate, sodium tripolyphosphate, sodium phosphate glass; neutral salts such as zeolites, sodium 10 sulfate, sodium chloride, and talc; silicates and silicate hydrates such as sodium metasilicate, sodium sequisilicate, sodium/potassium silicate water glasses, orthosilicate and mixtures thereof.

A useful initial inorganic coating compound for a halogen bleach core material to be used in a detergent composition would be an admixture of sodium sulfate and sodium tripolyphosphate. Sodium sulfate and sodium tripolyphosphate are relatively inert with respect to halogen bleaches and are components commonly blended into detergent compositions.

Outer Coating Materials

Nearly any substance may be employed as the outer coating material so long as it is solid at normal storage. temperatures (typically between 30° and 100° F.). 25 nonexhaustive list of compounds which may be used as the second coat includes alkalies such as sodium carbonate, sodium bicarbonate, sodium sequicarbonate, sodium borate, potassium bicarbonate, potassium sequicarbonate, potassium sodium sulfate hydrate, phosphates 30 diammonium phosphates, monocalcium phosphate, tricalcium phosphate, calcium pyrophosphate, iron pyrophosphate, magnesium phosphate, monopotassium orthophosphate, potassium pyrophosphate, disodium orthophosphate, trisodium orthophosphate, tetrasodium pyrophosphate, 35 tripolyphosphate, sodium phosphate glass; neutral salts such as zeolites, sodium sulface, sodium chloride, and talc; silicates and silicate hydrates such as sodium

metasilicate, sodium sequisilicate, dry sodium/potassium silicate water glasses, sodium orthosilicate; organic sequestering agents such as copolymers of vinylacetate and maleic anhydride, copolymers of acrylic acid and maleic anhydride, copolymers of maleic anhydride and itaconic acid, polyacrylic acid; and N-alkyl sulfonate, such as sodium carboxymethyl celluloses, octvl sulfonate, hydropropyl cellulose, hydroxyethyl ether of cellulose, hydroxypropylmethyl cellulose; C12 to C20 fatty acids such as stearic acid, palmitic acid, and n-alkanoic acids; paraffin waxes; microcrystalline waxes; C12 and greater primary and secondary solid alcohols; Pluronic surfactants with molecular weight between about 8,000 to about 16,500; primary and secondary alkyl sulfates; and alkali metal 15 sulfonates and mixtures thereof. The preferred encapsulate uses an outer coating that is insoluble in the liquid composition prior to casting. After the detersive system is cast, the preferred encapsulates are water soluble to permit release of the halogen into the cleaning medium.

The synthetic detergent compound used in the coating must remain sufficiently solid at temperatures likely to be encountered during storage of the product, for example, temperatures of about 15 to 50° C., and also remain stable at temperatures likely to be encountered during processing 25 of the product into end use mixtures, for example, temperatures of about 15 to 95° C.

20

Synthetic detergents that can be used include anionic, cationic, nonionic and amphoteric detergent compositions. Examples of anionic detergents useful in the detergent-30 bleach compositions of the invention are the higher alkyl mononuclear aromatic alkali-metal sulfates and sulfonates, alkyl sulfates and sulfonates linear alkylbenzene sulfonates having about 9 to about 13 carbon atoms in the alkyl group wherein the alkyl group is derived 35 from polypropylene as described by Lewis in U.S. Pat. No. 2,477,382, or wherein the alkyl group is a hexene dimer or trimer as in McEwan, U.S. Pat. No. 3,370,100, or wherein WO 90/12081 - 13 - PCT/US89/03313

the alkyl group is derived from alpha-olefins, as in Swenson, U.S. Pat. No. 3,214,462. Also there may be employed primary and secondary alkyl sulfates.

The soaps are included within the definition of anionic detergents as used herein. Examples of operable soaps soluble with the present ir /e. tion are the sodium and potassium salts of acyclic monocarboxylic acids having chain lengths of about 8 to about 22 carbon atoms.

A particularly suitable synthetic detergent for use as a coating in the present invention is preoxidized sodium octyl sulfonate. The sodium octyl sulfonate may contain 1,2 alkane bisulfonate as a by-product of manufacture which does not affect the performance of sodium octyl sulfonate as a coating in the invention.

The organic compound coating is applied as a solution in a suitable solvent, water being preferred because of its compatibility with chlorine releasing agents, non-flammability, and non-toxicity.

The compositions of the present invention may be 20 formulated with a detergent builder as a detergency aid, for example, those mentioned hereinafter, to provide a commercially valuable detergent-bleach composition.

Inorganic fillers suitable for coating agents include alkalies such as sodium bicarbonate, sodium sequicarbonate, 25 sodium borate, potassium bicarbonate, potassium sequicarbonate, potassium borate; phosphates such as diammonium phosphate, monocalcium phosphate monohydrate, tricalcium calcium pyrophosphate, iron pyrophosphate, phosphate, magnesium phosphate, monopotassium orthophosphate. 30 potassium pyrophosphate, disodium orthophosphate, dihydrate, trisodium orthophosphate, decahydrate, tetrasodium pyrophosphate, sodium tripolyphosphate, phosphate glass; neutral soluble salts such as sodium sulfate and sodium chloride; silicates; organic seques-35 tering agents; and anti-redeposition agents.

When carrying out the process of the instant invention, the protective encapsulate materials or coatings

of the invention are conveniently applied using fluidized bed encapsulating equipment. Such equipment comprises a coating chamber or cylindrical tower wherein the coating or encapsulation of the particles is accomplished. 5 unexpanded bed of the particles to be coated is introduced into the equipment. A nozzle constituting a spraying means is disposed within the equipment and adapted to be adjusted vertically so that the liquid coating material discharged in a downwardly diverging three-dimensional spray pattern 10 would just cover the upper surface area of the bed.

The coating solution is contained in a vessel and is fed to the nozzle. Fluidizing gas (typically air) passes into the fluidized bed area. The fluidized gas is heated or cooled if required, in order to maintain the fluidizing 15 gas within a desired temperature range.

A known weight of particles of a halogen source to be coated is placed in the equipment. Air is caused to flow into the fluidized bed thereby expanding the layer of particles, and maintaining the particles in continuous 20 motion within the volume defined by the expanded bed, thus forming a fluidized bed. A solution of a coating substance is sprayed through the nozzle on the fluidized bed of particles until all particles in the bed are completely coated. Particles coated by the above-described procedure 25 are completely encapsulated with a continuous coating, and are free-flowing a.d non-agglomerated.

It is important that each particle be fully covered to prevent the oxidizing halogen source from reacting in the detersive system environment.

30

When it is desired to apply an initial coating of a coating agent and a subsequent coating of a synthetic detergent, the double coating may be conducted in a single fluidized bed either by applying the first coat, emptying the solution tank, filling the solution tank with the 35 second coating solution and then applying the second coat; or with a dual coating solution inlet to the nozzle, the fluidized particles in the bed first being coated with the WO 90/12081 - 15 - PCT/US89/03313

coating agent contained in a solution tank, this first coating being allowed to dry and then a second coating of the synthetic detergent contained in a second solution tank being applied, both coatings being conducted in accordance with the previous discussion on the operation of the fluidized bed.

A third method of applying a double coating in a fluidized bed is to coat the core particles with the coating agent in a first fluidized bed apparatus. The coated material is then allowed to dry and placed in a second fluidized bed apparatus, wherein the encapsulated product produced in the first fluidized bed is coated with a second coating solution of a synthetic detergent. The fluidized bed operation is conducted in accordance with the prior discussion of the operation of the fluidized bed.

Before removal of the encapsulated oxidizing chlorine source from the fluidized bed the temperature in the bed can be increased so as to drive off any solvent remaining in the encapsulate. However, the temperature must remain below the melting temperature of the encapsulant and below the degradation temperature of the encapsulated core and coatings.

The encapsulated halogen bleach sources of the present invention comprise about 20 to 90 wt-% halogen bleach source core and about 10 to 80 wt-% coating when a single coating is utilized, and about 20 to 90 wt-% halogen bleach source core, about 0.5 to 50 wt-% inorganic coating agent first coat, and about 5 to 70 wt-% synthetic coating when a double coating is utilized.

More particularly, the single coated halogen bleach source comprises about 30 to 80 wt-% halogen bleach source core and about 20 to 70 wt-% synthetic coating and most particularly about 40 to 55 wt-% halogen bleach source core and 45 to 60 wt-% synthetic coating.

A more preferred embodiment of the double coated halogen bleach source comprises about 30 to 80 wt-% halogen bleach source core, about 5 to 50 wt-% inorganic coating

agent first coating, and about 5 to 50 wt-% synthetic coating. In a most preferred embodiment, the encapsulate comprises about 30 to 60 wt-% chlorine bleach source core, about 15 to 45 wt-% inorganic coating agent first coating, and about 10 to 35 wt-% detergent second coating.

Dispensing

The cast solid detersive systems of the invention can be dispensed from a manual or automatic dispenser in which a stream of water is contacted with a surface of the cast 10 material providing a concentrate that is directed to a use location.

The cast material can be contained in a disposable container and inserted into the dispenser in that form. Alternatively the cast material can be manually removed 15 from a disposable container directly into a dispenser wherein the stream of water is contacted with at least one surface of the material. Typical dispensers are mechanical apparatus containing a nozzle for directing a stream of water onto the solid cast material. The dispenser 20 typically comprises a housing containing the internal The housing typically includes a storage working parts. portion wherein the mass of the solid block detersive system can be supported. The dispenser contains typically a support means upon which the cast material is placed. 25 The support is typically horizontally mounted within the dispenser and maintains the block material in position adjacent to the spray. The preferred support comprises a screen mounted to the inner walls of the housing at a position fixed above the spray such that the spray contacts 30 the majority of the solid cast detersive system. spray-forming nozzle is connected to a pressurized source of water by means of a supply line. The spray controlled by a device that can demand the addition of the concentrate made by spraying water on the cast material. 35 Upon the receipt of demand, water flow is directed through the supply line and nozzle onto substantially the entire lower surface of the cast material at pressures typically WO 90/12081 - 17 - PCT/US89/03313

greater than 10 psi. Heated water can be used depending on the formulation. The use of heated water, all else being equal, increases the rate of dispensing. The detersive system passes in solution form through the support screen and is directed by underlying collector portion of the housing to an outlet port and through a conduit to a utilization point. The utilization point can be a warewashing machine, a station for charging containers such as buckets or other apparatus with a concentrate for dissolution with additional quantities of water, or other use positions.

Alternatively the dispensing apparatus can be manually operated such that a measured amount of concentrate can be produced by manually triggering the spray onto the cast 15 material.

We have found that the stability of the chlorine source in the presence of the organic compositions relates to the quantity of free water in the cast material. have found that the detersive systems during manufacture 20 require some water for processing. Preferably all water present in the detersive system after casting is in the form of bound water or water of hydration or complexed water which is sequestered and removed from reactivity with the components. Free water is water available for reaction 25 with the encapsulated chlorine material and can provide a medium of reaction between released chlorine and any organics present in the cast material. We have found that maintaining the concentration of free water below about 10 wt-% can aid in preserving the availability of chlorine in the cast material over a substantial period of time, 30 preferably maintaining the concentration of free water at less than 5 wt-% or most preferably less than 2 wt-% can be very effective in maintaining the stability of material.

35

Example I

Chlorinated General Purpose Cleaner

Into a stainless steel jacketed tank equipped with a

variable speed turbine agitator was charged 10 parts of soft water which was heated to 170° F. Into the mixed water was added 20 parts of a polyethylene glycol (CARBOWAX 8000, Union Carbide) at a sufficient rate to dissolve the 5 CARBOWAX in the water. Agitation was ended and into the tank was placed 22 parts of a linear alkyl sulfonate, 4 parts of a polyacrylic acid polymer (GOODRITE K-7058D, B. F. Goodrich), 14.5 parts of a powdered sodium tripolyphosphate, and 1 part of а granular 10 tripolyphosphate, 14 parts of anhydrous sodium metasilicate and 6 parts of sodium bicarbonate. The product was cooled to a temperature below 145° F. Next, 8.5 parts of an encapsulated sodium isocyanurate (of Example II) is added to the tank and the contents of the tank are carefully 15 agitated at a temperature below 145° F. in a manner such that the encapsulate is evenly distributed throughout the product. The material is drawn from the tank and 2 lbs. of is placed the warm liquid material in polyethylene The containers and their contents are cooled containers. in an air chiller for 12-15 minutes at a temperature less than 10° F. and are solidified.

Example II

20

The encapsulated sodium dichloroisocyanurate used in Example I was prepared as follows. Into a cylindrical 25 fluidized bed encapsulating machine was placed 61.25 parts sodium dichloroisocyanurate. The particles were fluidized and suspended in air by an upwardly moving stream of air at a temperature of about 90° C. Onto the heated suspended particulate was sprayed a solution comprising in 30 68.97 parts of soft deionized water, 16.5 parts of sodium sulfate and 5.3 parts of sodium tripolyphosphate. addition of this solution creates a first inorganic layer of mixed sodium sulfate and sodium tripolyphosphate. the complete addition of the inorganic layer, a solution 35 comprising 49.62 parts of a sodium linear alkyl sulfonate in 49.62 parts of soft water is then sprayed on the fluidized particle. This second step created an organic WO 90/12081 - 19 - PCT/US89/03313

encapsulate layer on the exterior of the particle. Substantially all free water (about 151.5 parts were removed) was removed during encapsulation.

Example III

chlorinated laundry detergent solid was according to the following procedure. Into a stainless steel mixing ta:k equipped with heating and cooling and a variable speed turbine agitator was placed 20 parts of a nonylphenol ethoxylate having an average of 9.5 mol.s of 10 ethylene oxide, and 0.6 part of hexylene glycol. The mixer is begun and into the tank is charged 11.35 parts of a polyethylene glycol (CARBOWAX 8000, Union Carbide) at a rate such that the polyethylene glycol melts and dissolves in the liquid phase. In a separate container 2.3 parts of soft water, 0.1 part of a fluorescent brightener and 0.004 part of a blue dye were mixed and then charged to the tank. The temperature of the charged materials was permitted to fall below 150° F. In a separate dry blender 0.5 part of carboxymethyl cellulose, 27.246 parts of 20 tripolyphosphate powder and 9.4 parts of anhydrous sodium metasilicate were dry blended. The dry blend was slowly added to the liquid phase in the mixing tank. uniformity was reached, 0.25 part of a fragrance was added. The mixture was cooled and to the caref lly agitated mixture was added 19 parts of a chlorinated isocyanurate manufactured in Example IV with 5 parts of sodium The agitation was controlled such that the carbonate. encapsulate was just distributed throughout the product and terminated. The material was packaged in 4 lb. aliquots in 30 polyethylene tubs and cooled until solid.

Example IV

Using the procedure of Example II the following encapsulate was prepared.

	Ingredient	Parts by Weight
	Core:	
	Sodium dichloroisocyanurate	63.2
	Coating 1:	
5	Sodium sulfate	25.8
	Sodium tripolyphosphate	8.6
	Soft water	103.0
	Coating 2:	
	Hydroxypropyl cellulose	4.9
10	Soft water	98.4
	Water removed during encapsulation	203.9

Example V

Using the procedure of Example II an encapsulated 15 chlorine source was prepared using the following ingredients.

	Ingredient	Parts by Weight
	Core:	
20	Sodium dichloroisocyanurate	80.65
	Coating 1:	
	Sodium sulfate	14.73
	Sodium tripolyphosphate	½.33
	Soft water	58.68
25	Coating 2:	
	Carboxymethyl cellulose	5.00
	Soft water	88.80
	Water removed during encapsulation	351.27

30 Example VI

Using the procedure of Example II an encapsulated chlorine source was prepared using the following ingredients.

	Ingredient	Parts by Weight
	Core:	
	Sodium dichloroisocyanurate	63.2
	Sodium sulfate	25.8
5	Coating 1:	
	Sodium tripolyphosphate	8.6
	Soft water	103.0
	Coating 2:	
	Hydroxyethyl cellulose	4.9
10	Soft water	98.3
	Water removed during encapsuation	203.9

Example VII

An encapsulated chlorine source was prepared using the 15 procedure of Example II using the following ingredients.

<u>Ingredient</u>	Parts by Weight
Core:	
Sodium dichloroisocyanurate	63.2
20 Coating 1:	
Sodium sulfate	25.8
Sodium tripolyphosphate	8.6
Soft water	103.0
Coating 2:	
25 Methyl cellulose	4.92
Soft water	245.92
Water removed during encapsulation	351.46

Example VIII

A solid chlorinated fabric softening laundry detergent was prepared following the procedure of Example III using the following ingredients.

	Ingredient	Parts by Weight
	Nonylphenol ethoxylate	26.39
	with 9.5 moles ethylene oxide	
	Hexylene glycol	1.17
5	Polyethylene glycol	14.66
	Soft water	2.25
	Dye	0.08
	Sodium tripolyphosphate	26.39
	Sodium metasilicate	3.81
10	Quaternary ammonium fabric softener	2.25
	Encapsulate of Example VI	23.00

Example IX

Example VI was repeated with the encapsulate of 15 Example II.

Example X

Example VIII was repeated with the encapsulate of Example $\mathbf{V}.$

Example XI

20 Example VIII was repeated with the encapsulate of Example VII.

Table 1

Active Chlorine Stability

25 <u>Percent Active Chlorine Retained After</u>

<u>Storage at 100° for Two Weeks</u>

	<u>Preparation</u>		Percent Retained
	Example	VIII	67.2
30	Example	IX	57.5
`	Example	X	62.8
	Example	XI .	85.2

Example XII

35 <u>Chlorinated General Purpose Cleaner</u>

Into a stainless steel mix tank equipped with heating and cooling equipment and a variable speed turbine type

WO 90/12081 - 23 - PCT/US89/03313

agitator was added 10 parts of water and 22 parts of polyethylene glycol (CARBOWAX 8000). The glycol was added at a rate such that it was melted and fully mixed upon addition. Into the heated solution was added 20 parts of a linear alkyl sulfonate, 4 parts of a polyacrylate polymer, 15.5 parts of sodium tripolyphosphate, 14.0 parts of sodium metasilicate, and 6 parts of sodium bicarbonate. The contents of the mixer were agitated until uniform and into the mixer was added 8.5 parts of the encapsulate of Example II. The contents of the mixer were carefully agitated until just uniform. The material manufactured above had a chlorine stability of 106.06% chlorine retained at 100° F. for two weeks of storage.

15 <u>Table 2</u>

<u>Dispensing Characteristics of Product of Example XII</u>

Grams Dispensed

		(Average of 5 Te	sts)
<u>Te</u>	emperature	At 30 psi	At 50 psi
20 12	20°	34.338	53.735
13	30°	38.72	59.2
14	00	50.68	
1 5	00		66.184
	, ,	55.44	81.7

25 The data in Table 2 snows that the product of Example XII is easily dispensed using warm water at commonly available temperatures at commonly available pressures at most end use sites. The product can easily be dispensed for any typical end use by controlling either pressure, temperature or dispensing time.

The product was dispensed by placing the material in a dispenser using a 30-second cycle at either 30 or 50 psi and at temperatures ranging from $120-150^{\circ}$ F.

A useful dispenser is shown in FIGURE 1. Referring to 35 Fig. 1, there is generally disclosed a dispenser having a container or housing 20. The housing has a generally cylindrical upper storage portion 21 having a cylindrical

inner wall 22. The wall 22 defines an internal cavity 23. The upper terminous of the storage portion 21 defines an access port 24 into cavity 23 of storage portion 21.

Inner wall 22 of housing 20 converges in the downward 5 direction, defining a lower funnel-shaped collector portion Inner wall 22 of housing 20 25 of housing 20. configured to form an annular flange at 26 circumferentially extending around inner wall 22 of housing 20 at the juncture of upper storage portion 21 and lower 10 collector portion 25. The lower terminous of collector portion 25 defines an outlet port 27 from internal cavity 23 for passage therethrough of solution collected by Outlet port 27 has a hose clamp collector portion 25. extension 28 having a plurality of annular ribs configured 15 for engaging the inner walls of a connecting hose or conduit 29.

The outlet port 27 may be directly connected with the wash chemical solution utilization point by conduit 29 and feed thereto by gravity as it is created or feed thereto by 20 a wash chemical solution pump 30 placed in conduit 29.

Housing 20 may be constructed of any suitable material which is capable of withstanding exposure to cleaning solutions, and is preferably configured of stainless steel molded plastic material. The housing 20 can 25 constructed of a transparent or translucent material to allow the operator to see at a glance the amount of wash chemical in storage portion 21 and if dispenser 20 neess to be refilled. If housing 20 is not made of a transparent or translucent material, preferably a portion of storage 30 portion 21 is made transparent or translucent to aid in determining when dispenser 20 should be refilled. 32 are connected to mounting plates and extend rearwardly from the outer surface of housing 20 securely mounting housing 20 to a vertical side wall.

A door 34 is sized to extend entirely across and to sealingly close access port 24. Door 34 is pivotally mounted at 35 for pivotal motion between a closed and

35

WO 90/12081 - 25 - PCT/US89/03313

opened position. The lower collector portion 25 of housing 20 has an outwardly projecting coupling portion 36 extending from collector portion 25 adjacent outlet port 27 of collector portion 25. A tube fitting insert 37 is secured within coupling projection 36 and projects through inner wall 22 of collector portion 25 of housing 20. A spray-forming nozzle 38 is threaded into the end of tube insert 37 and is axially aligned within inner cavity 23 of housing 20 in a direction so as to direct an upwardly projected spray pattern therefrom. Tube fitting insert 37 is provided with an O-ring seal 39.

10

A horizontal support screen 40 is mounted in resting engagement upon annular flanged portion 26 of housing 20. Support screen 40 nas about 1 inch square openings in order to support a solid block of detersive system 80 without significantly interfering with the impingement of water sprayed from nozzle 38 onto the lower surface of water sprayed from nozzle 38 onto the lower surface 81 of the detersive system 80 (i.e., the surface in contact with support screen 40).

A 1/4 to 1/20 inch (0.63 to 0.13 cm) lower screen 41 is placed in collector portion 25 of housing 20 between spray nozzle 38 and outlet port 27 to catch any undissolved chunks of wash chemical 80 small enough to pass through support screen 40. This prevents small chunks of chemical 80 collecting in outlet port 27 or conduit 29 and blocking the flow of concentrated wash chemical solution out of dispenser 20.

A water supply inlet pipe 42 is connected to tube

insert 37 and is in communication therewith for providing a source of water flow to spray-forming nozzle 38. Water supply line 42 passes through one of the mounting plate members 32 and receives structural support therefrom. A siphon breaker 43 interrupts water supply line 42. A

safety switch 50 is mounted to door 34 for movement therewith and senses the operative position of door 34 relative to access port 24 of housing 20. In the preferred

embodiment, safety switch 50 comprises a mercury actuated switch.

FIGURES 2 and 3 represent an embodiment of the product format for the cast detersive systems of the invention.

5 The solid cast detersive system of the invention is packaged in a closable container 200 which comprises a lid 210 and a lower container section 211. The lid 210 can be made of a variety of materials including paper, film, foil, etc. The lower container can also be made of a variety of materials, however thermoplastic deformable material is preferred. The lid 210 is adhered to the container 211 through a releasable layer 212 disposed between the lid and the container.

FIGURE 3 shows a side view of the cast detersive system within its container. In use lid 210 is preled from the top of the lower portion revealing the upper surface of the cast material 80. The container as a whole without the lid can be inserted into the dispenser. Alternatively the container body 211 if made of a deformable material can be flexed for the purpose of removing the cast material 80 in a solid block which can then directly be inserted into the dispenser for contact with a water stream.

While the foregoing discussion, Examples and data provide a basis to understand the invention, many embodiments of the invention can be made without departing from the spirit and scope of the invention. Thus the invention resides in the claims hereinafter appended.

WO 90/12081 - 27 - PCT/US89/03313

WHAT IS CLAIMED IS:

35

- 1. A solid cast, active halogen-containing,
 storage-stable, detersive system comprising:
- (a) an effective detersive amount of a source of alkalinity;
 - (b) an effective amount of a hardness sequestering agent;
 - (c) about 5 to 60 wt-% of an encapsulated source of active halogen; and
- 10 (d) about 5 to 50 wt-% of a polyalkylene oxide polymer;

wherein, in the cast composition, the halogen available from the source of active halogen is storage stable in the presence of the polyalkylene oxide polymeric composition.

- 15 2. The detersive system of claim 1 wherein the halogen is chlorine.
 - 3. The detersive system of claim 2 wherein there is 15 to 30 wt-% of the encapsulated source of active chlorine.
- 20 4. The detersive system of claim 2 wherein there is 15 to 50 wt-% of the encapsulated source of active chlorine.
- 5. The detersive system of claim 2 wherein there is: 20 to 35 wt-3 of the source of encapsulated active 25 chlorine.
 - 6. The detersive system of claim 2 wherein the encapsulate comprises 1 to 80~wt-% of a source of active chlorine and 99 to 20 wt-% of an encapsulating layer.
- 7. The detersive system of claim 6 wherein the 30 encapsulating layer comprises a synthetic anionic surfactant.
 - 8. The detersive system of claim 6 wherein the encapsulating layer comprises 10 to 90 wt-% of a first inorganic layer and 90 to 10 wt-% of a second organic layer.
 - 9. The detersive system of claim 8 wherein the organic encapsulating layer comprises a cellulosic layer, a

synthetic anionic surfactant or mixtures thereof.

- 10. The detersive system of claim 9 wherein the cellulosic layer comprises methyl cellulose, a hydroxyalkyl cellulose, or mixtures thereof.
- 11. The detersive system of claim 2 wherein the source of active chlorine is selected from the group trichloroisocyanuric acid, consisting of potassium dichloroisocyanurate, sodium dichloroisocyanurate, sodium dichloroisocyanurate dihydrate, and mixtures thereof.
- The detersive system of claim 1 wherein the 10 polymeric composition comprises a polyalkylene oxide polyethylene oxide polymer, a polypropylene oxide polymer, or an ethylene oxide/propylene oxide block copolymer.
- 13. The detersive system of claim 12 wherein the alkylene oxide polymer comprises a nonionic surfactant.
 - 14. The detersive system of claim 13 wherein the nonionic surfactant comprises a block copolymer containing at least one ethylene oxide block and at least propylene oxide block.
- The detersive system of claim 14 wherein the 20 15. nonionic surfactant comprises an alkyl phenol alkoxylate wherein the alkyl group is a $C_{1-1,2}$ alkyl and the alkoxylate comprises 2 to 24 moles of ethylene oxide.
- The detersive system of claim 1 wherein the 25 hardness sequestering agent comprises an organic hardness sequestering agent or an inorganic hardness sequestering agent.
- The detersive system of claim 16 wherein the organic hardness sequestering agent comprises a polyacrylic 30 acid, an organic phosphonate, or mixtures thereof.
 - The detersive system of claim 16 wherein the inorganic hardness sequestering agent comprises a condensed phosphate hardness sequestering agent.
- 19. The detersive system of claim 18 wherein the 35 condensed phosphate comprises alkali an tripolyphosphate sequestering agent.
 - The detersive system of claim 1 wherein the

source of alkalinity comprises an alkali metal hydroxide or an alkali metal silicate having an ${\rm M_2O:SiO_2}$ ratio of about 2:1 to 1:3.6 wherein M is an alkali metal.

- 21. The detersive system of claim 1 wherein the 5 source of alkalinity comprises an alkali metal carbonate, an alkali metal bicarbonate, an alkali metal borate and mixtures thereof.
 - 22. The detersive system of claim 1 which additionally comprises an anionic surfactant.
- 23. The detersive system of claim 22 wherein the anionic surfactant comprises an alkyl sulfonate, an alkyl sulfate, an alkyl benzene sulfate, or mixtures thereof.
- 24. A solid, cast, stable, chlorine-containing laundry detergent comprising:
 - (a) about 0.1 to 50 wt-% of a soil-removing nonionic surfactant;
 - (b) about 0.1 to 95 wt-% of a source of alkalinity;
- (c) an effective amount of a hardness sequestering agent;
 - (d) about 5 to 60 wt-% of an encapsulated chlorinated isocyanurate compound; and
- (e) about 5 to 50 wt-% of a polyalkylene glycol 25 casting agent;
 - wherein, in the cast laundry detergent composition, the available chlorine is storage stable in the presence of the nonionic surfactant and the polyalkylene glycol casting agent.
- 30 25. The detergent of claim 24 wherein there is 15 to 30 wt-% of the encapsulated source of active chlorine.
 - 26. The detergent of claim 24 wherein there is 20 to 25 wt-% of the source of encapsulated active chlorine.
- 27. The detergent of claim 24 wherein the encapsulate comprises 1 to 80 wt-% of a source of active chlorine and 99 to 20 wt-% of an encapsulating layer.
 - 2). The detergent of claim 27 wherein the

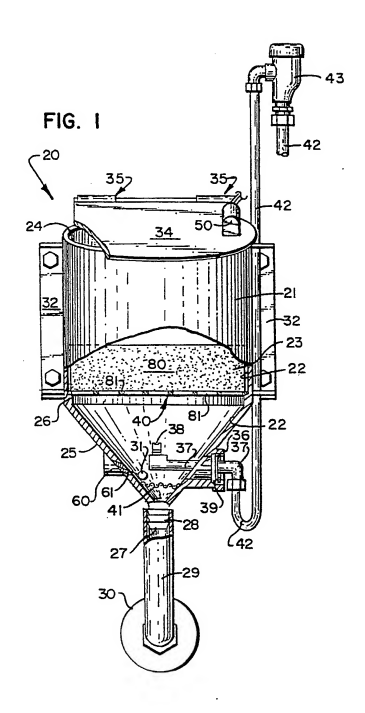
encapsulating layer comprises a synthetic anionic surfactant.

- 29. The detergent of claim 27 wherein the encapsulating layer comprises 10 to 90 wt-% of a first inorganic layer and 90 to 10 wt-% of a second layer comprising an organic encapsulating layer.
 - 30. The detergent of claim 29 wherein the organic encapsulating layer comprises a cellulosic layer, a synthetic anionic surfactant or mixtures thereof.
- 10 31. The detergent of claim 30 wherein the cellulosic layer comprises methyl cellulose, a hydroxyalkyl cellulose, or mixtures thereof.
- 32. The laundry detergent of claim 24 wherein the detergent additionally comprises an effective amount of a 15 soil anti-redeposition agent.
 - 33. The detergent of claim 32 wherein the soil antiredeposition agent comprises a carboxymethyl cellulose composition.
- 34. The detergent of claim 24 wherein the detergent 20 additionally comprises a dye, a perfume, an anti-redeposition agent, a brightener or mixtures thereof.
 - 35. The detergent of claim 24 wherein the chlorinated isocyanurate compound is selected from the group consisting of trichloroisocyanuric acid, potassium dichloroisocyanurate, sodium dichloro isocyanurate, sodium dichloroisocyanurate dihydrate, and mixtures thereof.
- 36. The detergent of claim 35 wherein the nonionic surfactant comprises a block copolymer containing at least one ethylene oxide block and at least one propylene oxide 30 block.
 - 37. The detergent of claim 36 wherein the nonionic surfactant comprises an alkyl phenol alkoxylate wherein the alkyl group is a C_{1-12} alkyl and the alkoxylate comprises 2 to 24 moles of ethylene oxide.
- 35 38. The detergent of claim 24 wherein the hardness sequestering agent comprises an organic hardness sequestering agent or an inorganic hardness sequestering

WO 90/12081 - 31 - PCT/US89/03313

agent.

- 39. The detergent of claim 38 wherein the organic hardness sequestering agent comprises a polyacrylic acid, an organic phosphonate, or mixtures thereof.
- 40. The detergent of claim 38 wherein the inorganic hardness sequestering agent comprises a condensed phosphate hardness sequestering agent.
- 41. The detergent of claim 40 wherein the condensed phosphate comprises an alkali metal tripolyphosphate 10 sequestering agent, as alkali metal carbonate, an alkyl metal bicarbonate, and mixtures thereof.
- 42. The detergent of claim 24 wherein the source of alkalinity comprises an alkali metal hydroxide or an alkali metal silicate having an M₂O:SiO₂ ratio of about 2:1 to 1:3.6 wherein M is an alkali metal.
 - 43. The detergent of claim 24 which additionally comprises an anionic surfactant.
 - 44. The detergent of claim 24 which additionally comprises a quaternary fabric softener.



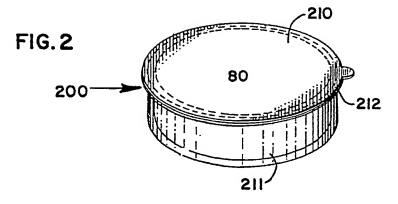
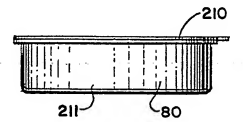


FIG.3



INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 89/03313

I. CLASS	IFICATION OF SUBJECT MATTER (If several classification symb	ols apply, indicate all) 6
According	to International Patent Classification (IPC) or to both National Classific	ation and IPC
	C 11 D 3/395, C 11 D 17/00, C 13	
II. FIELD	SEARCHED	
	Minimum Documentation Search	ned ?
Classification	on System Classification	Symbols
IPC ⁵	C 11 D	
	Documentation Searched other than Minimum to the Extent that such Documents are Included	
III. DOCL	IMENTS CONSIDERED TO BE RELEVANT	
Category *	the state of the s	e relevant passages 12 Relevant to Claim No. 13
х	EP, A, 0307587 (ECOLAB INC.) 22 March 1989 see page 6, lines 13-45; and claims	1-43
A	EP, A, 0003769 (ECONOMICS LAB. 5 September 1979 see claims	1
A	EP, A, 0264043 (HENKEL) 20 April 1988 see claims	1
. A	US, A, 3637509 (W.R. BRENNAN) 25 January 1972 see claims	1
A	US, A, 4136052 (L.R. MAZZOLA) 23 January 1979 see claims	1
A	EP, A, 0298222 (ECOLAB INC.) 11 January 1989	1
"A" doc cor "E" eer filir "L" doc wh cute "O" doc oth "P" doc late	cument defining the general state of the art which is not cited invertised end to be of particular relevance. Iter document but published on or after the international grate cument which may throw doubts on priority claim(s) or ch is cited to establish the publication date of another claim or other special reason (as specified) "Y" document referring to an oral disclosure, use, exhibition or er means cument published prior to the international filing date but in the priority date claimed "IFICATION	document published after the international filing date tority date and not in conflict with the application but to understand the principle or theory underlying the stion ment of particular relevance; the claimed invention of be considered novel or cannot be considered to ve an inventive step ment of particular relevance; the claimed invention of be considered to involve an inventive step when the ment is combined with one or more other such docus, such combination being obvious to a person skilled e art. The property of the same patent family
I	h October 1989	1 0 NOV.1989
internation	EUROPEAN PATENT OFFICE	T.K. WILLIS

ĩ

Category * 1	NTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHE Citation of Document, with indication, where appropriate, of the relevant passages	i Relevent to Claim h
	see claims	
Ì		
i		
:		
į		
į		
! !		
i		
1		
-		<u> </u>
;		
İ		
:		
		•
•		
•		
i		
	•	
		•
i		

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 8903313

SA 30516

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 02/11/89
The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date		Patent family member(s)	
EP-A- 0307587	22-03-89	AU-A-	1461988	16-02-89
EP-A- 0003769	05-09-79	BE-A- CA-A- JP-A- US-A- US-A- US-E- US-E-	904138 1125621 54152007 4569781 4569780 RE32763 RE32818	15-05-86 15-06-82 29-11-79 11-02-86 11-02-86 11-10-88 03-01-89
EP-A- 0264043	20-04-88	DE-A-	3634812	14-04-88
US-A- 3637509	25-01-72	None		
US-A- 4136052	23-01-79	US-A- AU-B- AU-A- CA-A- GB-A- JP-A- US-A- US-A-	4078099 508961 2806777 1095662 1591335 53026782 4327151 4126717	07-03-78 17-04-80 20-09-79 17-02-81 17-06-81 13-03-78 27-04-82 21-11-78
EP-A- 0298222	11-01-89	US-A- JP-A-	4830773 1031899	16-05-89 02-02-89